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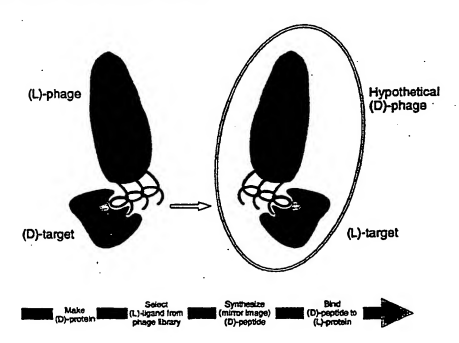
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(57) Abstract

A method of identifying macromolecules (peptides, olignucleotides, sugars and macromolecular complexes, such as RNA-protein complexes, protein-lipid complexes), which are not of the natural handedness (not of the chirality as they occur in nature or as a wild type molecule) and which are lighds for other chiral macromolecules.

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IDENTIFICATION OF ENANTIOMERIC LIGANDS

Related Applications

This application is a Continuation-in-part of U.S. Serial Number 08/627,497, filed March 28, 1996, which is a 5 continuation-in-part of U.S. provisional application number 60/001,067, filed July 11, 1995, and is a continuation-inpart of U.S.S.N. 08/482,309, filed June 7, 1995, which is a continuation-in-part of U.S.S.N. 08/433,572, filed May 3, 1995, the teachings of which are incorporated herein by reference.

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Background

Genetically encoded libraries of peptides and 15 oligonucleotides are well suited for the identification of ligands for many macromolecules. However, a major drawback of biologically encoded libraries is that the resultant ligands are subject to degradation by naturally occurring 20 enzymes. Furthermore, because of their sensitivity to cellular proteases, peptides composed of naturally occurring L-amino acids are efficiently processed for major histocompatibility complex class II-restricted presentation to T helper cells (T_H cells). As a result, L-peptides can induce a vigorous humoral immune response that impairs the activity of such drugs (Gill, T.J., et al., Nature, 197:746 (1963); Mauer, J., J. Exp. Med., 121:339 (1965); Borek, F., et al., Biochem. J., 96:577 (1965); Janeway, C.A. and Sela, M., Immunol., 13:29 (1967); Dintzis, H.M., et al., Proteins, 16:306 (1993)).

The enantiomers of macromolecules of natural handedness make better drugs than the macromolecules of natural handedness. In contrast to naturally occurring L-peptide sequences and D-nucleic acid sequences, the enantiomers of these naturally occurring macromolecules (e.g., D-peptides and L-nucleic acids) are not good substrates for naturally occurring proteases and nucleases. In addition, the enantiomers of naturally occurring molecules do not elicit an efficient immune response.

Availability of D-peptides and L-nucleic acids for use as drugs is desirable.

Summary of the Invention

The present invention is a method of identifying enantiomeric macromolecules (proteins, peptides, 15 oligonucleotides, nucleic acids, sugars and macromolecular complexes, such as RNA-protein complexes and protein-lipid. complexes), which are not of the natural handedness (not of the chirality as they occur in nature or as a wild type molecule) and which are ligands for other chiral macromolecules, which are referred to as target or desired 20 macromolecules. Target or desired macromolecules include oligonucleotides (DNA, RNA) and proteins (e.g., polypeptides and peptides), such as hormones, enzymes, antibodies and antigens. In one embodiment, the present invention is a method of identifying D-amino acid peptide ligands which bind a target or desired L amino acid peptide. In a second embodiment, this invention is a method of identifying peptides comprised of D-amino acid residues that are ligands for oligonucleotides (RNA or DNA). In a further embodiment, the present invention is a method of identifying RNA or DNA oligonucleotides which are

of the opposite chirality from that which occurs in nature. DNA occurs in nature as a D isomer.

In one embodiment, the present invention relates to a method of producing a macromolecule of non-natural 5 handedness that binds to a target macromolecule of natural handedness (e.g., peptide, oligonucleotide), which is performed as follows: an enantiomer of the target macromolecule or of a domain characteristic of the target molecule is provided and contacted with a library of macromolecules of natural handedness, under conditions 10 appropriate for binding of a macromolecule of natural handedness in the library with the enantiomer; as a result, the enantiomer binds a macromolecule of natural handedness present in the library. The enantiomer of the 15 macromolecule of natural handedness which is bound to the enantiomer of the target macromolecule is produced; the enantiomer of the macromolecule of natural handedness is a macromolecule of non-natural handedness which binds to the target macromolecule of natural handedness. That is, the 20 enantiomer of a macromolecule which is present in the library and binds to the enantiomer of the target molecule is produced; the result is a macromolecule of non-natural handedness which binds the target molecule (of natural handedness).

In the embodiment where the target macromolecule is a protein (e.g., peptide), a D amino acid peptide that binds to a target L peptide is produced. The method is performed as follows: a D amino acid peptide of the target L macromolecule or of a domain characteristic thereof and a 30 library of L amino acid peptides are provided. The library and the D amino acid peptide of the target macromolecule are contacted under conditions appropriate for binding of an L amino acid peptide in the library with the D amino acid peptide; as a result, the D amino acid peptide binds 35 an L amino acid peptide present in the library. An L amino WO 96/34879 PCT/US96/06155

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acid peptide present in the library which is bound to the D
amino acid peptide is identified and sequenced. A D amino
acid peptide of the L amino acid peptide identified in the
library or of a characteristic domain thereof is produced;
the resulting D amino acid peptide binds to the target L
macromolecule of natural handedness. That is, the
enantiomer of the L amino acid peptide which is present in
the library and binds to the D amino acid peptide of the
target L macromolecule is produced; the result is a D amino
acid peptide which binds the L-target macromolecule, which
in this embodiment is an L amino acid peptide.

In the embodiment, where the target macromolecule is a protein (e.g., peptide), an L oligonucleotide that binds to a target L protein is produced. The method is performed as 15 follows: a D amino acid peptide of the target L macromolecule or of a domain characteristic thereof and a library of D oligonucleotides are provided. The library is contacted with the D amino acid peptide under conditions appropriate for binding of a D oligonucleotides in the library with the D amino acid peptide, whereby the peptide 20 binds a D oligonucleotides present in the library. oligonucleotide which is bound to the D amino acid peptide is identified and sequenced. An L oligonucleotide of the D oligonucleotide identified in the library or of a 25 characteristic domain thereof, is produced; the L oligonucleotide binds to the target L macromolecule of nonnatural handedness. That is, the enantiomer of the D oligonucleotide which is present in the library and binds to the D amino acid peptide of the target L macromolecule 30 is produced; the result is an L oligonucleotide which binds the L-target macromolecule, which in this embodiment is an L amino acid peptide.

In another embodiment, the present invention relates to a method of producing a macromolecule of non-natural handedness that binds to a target macromolecule of natural

handedness, which is performed as follows: an enantiomer of the target macromolecule or of a domain characteristic of the target molecule is provided and contacted with a library of macromolecules of natural handedness, under 5 conditions appropriate for binding of a macromolecule of natural handedness in the library with the enantiomer; as a result, the enantiomer binds a macromolecule of natural handedness present in the library. A macromolecule of natural handedness which is bound to the enantiomer is 10 identified and sequenced. The enantiomer of the macromolecule of non-natural handedness, which is bound to the enantiomer of the macromolecule of natural handedness or of a characteristic domain thereof, is produced; the resulting enantiomer of the macromolecule of natural 15 handedness is a macromolecule of non-natural handedness which binds to the target macromolecule of natural handedness.

In a further embodiment of the present invention, an L
amino acid peptide which binds a D amino acid peptide of
interest is identified as follows: a phage display library
which comprises L amino acid peptides displayed on phage
surfaces is provided and contacted with the D amino acid
peptide of interest, under conditions appropriate for
binding of L amino acid peptides displayed on phage

25 surfaces with the D amino acid peptide of interest. Phage
which have on their surfaces the D amino acid peptide of
interest, bound to an L amino acid peptide displayed on the
surface (i.e., which have on their surfaces a D amino acid
peptide-displayed L amino acid peptide complex) are

30 identified. The displayed L amino acid peptide in the
complex is an L amino acid peptide which binds the D amino
acid of interest.

Optionally, the amino acid sequence of the L amino acid peptide displayed on the surface of the phage can be determined and the D amino acid peptide which corresponds

to the amino acid sequence of the L amino acid peptide can be synthesized, resulting in production of a D amino acid peptide which corresponds to the L amino acid peptide displayed on the phage surface. In one embodiment, 5 described herein, L amino acid peptides displayed on phage surfaces bind D amino acid peptides of a class of proteins, specifically the SRC homology 3 domain (SH3 domain).

A further embodiment is a method of making a D amino acid protein which corresponds to a target L amino acid 10 protein, which can be any protein (including polypeptides, peptides) for which a binding peptide is desired. In this embodiment, a phage display library which comprises a mixture of proteins displayed on phage surfaces is contacted with a D amino acid peptide corresponding to the 15 target L amino acid protein or corresponding to a domain characteristic of the target L amino acid protein, under conditions appropriate for binding of L amino acid peptides displayed on phage surfaces with D amino acid proteins. The mixture comprises the target L amino acid protein or a 20 characteristic L amino acid peptide domain thereof. Phage which have on their surfaces the D amino acid peptide bound to an L amino acid peptide displayed on the surface are identified. The amino acid sequences of the L amino acid peptides displayed on the surfaces of phages identified are 25 determined and a D amino acid protein which corresponds to an amino acid sequence of an L amino acid peptide is synthesized, resulting in production of a D amino acid peptide which corresponds to the target L amino acid protein.

The present invention also relates to a method of obtaining an L oligonucleotide nucleic acid sequence which binds an L amino acid peptide of interest. In this method, a collection of D nucleic acid sequences (e.g., a DNA library) is provided and contacted with a D amino acid 35 peptide of interest, under conditions appropriate for

binding of the D nucleic acid with the D amino acid peptide of interest. A D nucleic acid which binds to the D amino acid peptide is isolated and the nucleotide sequence of the D nucleic acid is determined. The D nucleic acid sequence 5 which binds to the D amino acid peptide of interest is prepared using L nucleotides, resulting in the production of an L nucleic acid sequence which binds an L amino acid peptide. A further embodiment of the invention relates to a method of obtaining an L nucleic acid sequence which 10 binds a D nucleic acid which comprises providing a collection of D nucleic acid sequences and contacting the D nucleic acid sequences with an L nucleic acid sequence. A D nucleic acid sequence which binds to the L nucleic acid sequence, thereby producing a D nucleic acid sequence - L 15 nucleic acid sequence complex, is identified. nucleotide sequence of the D nucleic acid sequence which binds to the L nucleic acid sequence is determined. nucleic acid sequence is synthesized using L nucleotides, resulting in the production of an L nucleic acid sequence 20 which binds a D nucleic acid.

Also the subject of the present invention are synthetic D amino acid peptides, such as D amino acid peptides identified and produced by the methods described herein, including but not limited to synthetic amino acid peptides which bind the SH3 domain, synthetic D amino acid peptides corresponding to all or a portion of the SH3 domain and, more generally, synthetic D amino acid peptides which bind a domain of an intracellular signaling protein. In addition, oligonucleotides (RNA, DNA) of non-natural handedness, such as oligonucleotides identified and produced by the methods described herein are the subject of this invention.

The present invention also relates to a process for producing a derivative of a macromolecule of non-natural handedness that binds a target macromolecule of natural

handedness. The method comprises the steps of identifying the macromolecule of non-natural handedness using the methods described herein and modifying or derivatizing the macromolecule of non-natural handedness to produce a derivative thereof. Derivatives obtainable by the method of the present invention described herein are also encompassed by the present invention.

D amino acid peptides and L nucleic acid sequences of the present invention are useful as drugs. For example, D amino acid peptides are not good substrates for naturally-occurring proteases (i.e., resistant to proteolytic degradation) and do not elicit an immune response comparable to that elicited by L amino acid peptides.

Brief Description of the Figures

The Figure is a graphic representation of the identification of a D-peptide ligand through mirror-image phage display.

Detailed Description of the Invention

The synthetic enantiomer of a structured biopolymer

folds into the mirror-image conformation of the natural

molecule; likewise, for a bimolecular complex, the two
enantiomers of the original partner molecules also form a
complex, with mirror-image symmetry to the original. The
present invention is based on the discovery that

identification of a macromolecule of natural handedness
(e.g., L-peptide, D-single-stranded oligonucleotide) that
binds the enantiomer of a chiral biological target
molecule, provides for a method of identifying a
macromolecule of non-natural handedness which binds the
natural form of the target. Such enantiomeric
macromolecules are assessed to determine their ability to
interfere with the biological activity of the target.

Thus, the present invention provides an approach to the development of new long-acting therapeutic or diagnostic molecules.

The present invention relates to a method of 5 identifying enantiomeric macromolecules, including peptides, polypeptides, proteins, oligonucleotides and sugars, as well as macromolecular complexes (oligonucleotide-protein complexes, protein-lipid complexes), which are not of the naturally-occurring or 10 wildtype handedness (i.e., chirality) and which are ligands for other chiral molecules (peptides, oligonucleotides and macromolecular complexes). As defined herein, an enantiomer of a macromolecule of natural handedness is the equivalent of the macromolecule of natural handedness, but 15 is of non-natural handedness. In the method of the present invention, an enantiomer of a naturally occurring target macromolecule is prepared and used to isolate, from a collection of naturally occurring macromolecules, a naturally occurring ligand that interacts with the 20 enantiomer. The enantiomeric form of the isolated naturally occurring ligand will interact with (e.g., bind) the naturally-occurring target macromolecule.

The target macromolecule of natural handedness can be any macromolecule having one or more chiral centers. The target macromolecule (chiral targets) can be intracellular or extracellular and includes, but is not limited to, nucleic acids (DNA, RNA), proteins or a characteristic domain thereof (e.g., peptide), peptides, polypeptides, oligonucleotides, carbohydrates, sugars, oligonucleotide-protein complexes (RNA-protein complex) protein-lipid complexes, and phospholipids, all of which contain chiral centers. Domains, fragments of regions of these macromolecules target macromolecules. The target macromolecules can be of mammalian origin (e.g., human) or non-mammalian origin (e.g., bacterial, fungal, viral, protozoan).

Examples of target macromolecules which are proteins (polypeptides, peptides) include a intracellular signaling proteins and domains thereof (e.g., the SH3 domains, the SH2 domains, the PH domains) (see Cohen, G.B., et al., 5 Cell, 80:237-248 (1995)), chemokines (e.g., α -chemokine, β chemokine) (See Clore, M. G., et al., FASEB J., 9:57-62 (1995)), cytokines (e.g., IL-1, TNF, lymphotoxin- α , IL-1 β , IL-6, M-CSF, TGFα), enzymes (e.g., protein kinase C, phospholipase C, phospholipase D) (See Divecha, N. and Irvine, R. F., Cell, 80:269-278 (1995)), tyrosine kinases (See Marshall, C. J., Cell, 80:179-185 (1995)), polypeptide growth factors, and domains thereof, the growth factor receptors and domains or fragments thereof (e.g., growth factors and growth factor receptors in the PDGF family, EGF 15 family, FGF family, IGF family, HFG family, VEGF family, neurotrophin family, Eph family, Class I cytokine family, GH family, IL-3 family, IL-6 family, IL-2 family, Class II cytokine family, TNF family) (See Heldin, C-H., Cell, 80:213-223 (1995), protein kinases, protein phosphatases, 20 cyclins and Cdc proteins (See Hunter, T., Cell, 80:225-236 (1995)), transcription factors and domains thereof (e.g., Ets domain, bZIP, rel homology domain, STATs, NF-ATs, TCF, Fos, JAKs) (See Hill, C.S. and Treisman, R., Cell, 80:199-211 (1995)) and hormones. Other examples of target 25 macromolecules for use in the present invention include: human CD2, human CD58 (LFA-3), human endothelin, heregulin- α , human interleukin-1 β converting enzyme (ICE), human macrophage inflammatory protein $1-\beta$, platet factor 4, human melanoma growth stimulating activity, GRO/melanoma growth 30 stimulating activity, MHC molecules, bacterial muramidase, kringle domains (e.g., plasminogen, apolipoproteins), ras, ras-GAP, selections (e.g., E-selectin, L-selectin, Pselectin), Pleckstrin homology domains, stromelysin, thrombin, tissue factor, calmodulin, CD4, collagenase, 35 dihydrofolate reductase, fibronectin, fibronectin type III

modules, G-protein subunits, vasopressin, Factor IX GLA domain, interleukin-8 (Clore, G.M., et al., Biochemistry, 29:1689-1696 (1990)), thrombomodulin EGF-like domain (Lentz, S.R., et al., J. of Biological Chem.,

5 288(20):15312-15317 (1993)), GPII_b-III_a cytoplasmic domain (Muir, T.W., et al., Biochemistry, 33:7701-7708 (1994)), Factor VIIa GLA domain, Factor IX EGF-like domain (Yang, Y., Protein Science, 3:1267-1275 (1994)), human immunodeficiency virus (HIV) proteins (e.g., HIV protease,

integrase, matrix, protein tyrosine phosphatase, reverse transcriptase, nef, tat, rev, envelope, and other HIV proteins, domains, fragments or scaffold mimics thereof), NH2-terminal SH3 domain GRB2, (Wittekind, M., et al., Biochemistry, 33:13531-13539 (1994)), COOH-terminal SH3

15 domain GRB2 (Kohda, D., et al., Structure, 2:1029-1040 (1994), P120^{GAP} SH3 domain (Yang, Y.S., et al., The EMBO Journal, 13(6):1270-1279 (1984)), and vascular permeability factor and vascular endothelial growth factor.

Examples of target macromolecules which are

20 oligonucleotides (RNA, DNA) include HIV RRE (rev responsive element), HIV Tar, and BCR-AB1 fusion DNA sequences.

Examples of target macromolecules which are phospholipids include phosphoinositide, phosphoinositidase C, phosphoinositide 3-kinase, phosphatidylinositol,

- phosphatidylinositol 3-phosphate, phosphatidylinositol
 (4,5)bisphosphate, phosphatidylinositol
 (3,4,5)triphosphate, phosphatidylcholine,
 phosphatidylethanolamine, phosphatidic acid, inositol (1,4)
 bisphosphate, inositol (1,4,5) triphosphate,
- diacylglycerol, sphingosine, sphingosine phosphate, sphigosine phosphocholine and ceramide (See Divecha, N. and Irvine, R. F., Cell, 80:269-278 (1995)).

In the method of the present invention, an enantiomer of a naturally occurring macromolecule (e.g., the

35 enantiomer of the target macromolecule or the enantiomer of

the macromolecule of natural handedness identified in the library) is prepared using routine methods. The enantiomer of the naturally occurring macromolecules can be prepared through the use of components of the opposite handedness from that which occurs in nature (e.g., the use of D amino acids for synthesis of mirror image peptides or the use of L-nucleic acids for synthesis of mirror image oligonucleotides). For example, a D-peptide for use in the present invention can be synthesized chemically and purified using affinity chromatography, as described in Example 1. Other methods for preparing the enantiomers of naturally occurring macromolecules are known in the art.

The full enantiomeric form of the target macromolecule or part of the three-dimensional molecular surface of the 15 target macromolecule can be used in the methods of the present invention. For example, the enantiomeric form of a subdomain of the target macromolecule, that by itself attains a conformation that resembles that of the enantiomeric form of this domain in the full macromolecule, 20 can be prepared and used in the methods of the present invention (Schumacher, T., et al., Science, 271:1854-1857 (1996)). Alternatively, continuous or discontinuous fragments of the enantiomeric form of the target macromolecule can be prepared on a peptidic or non-peptidic scaffold in which the composite surface of these fragments resembles part of the molecular surface of the full enantiomeric form of the target macromolecule (Tolman, R.L., et al., Int. J. Pept. Prot. Res., 41:455 (1993); Muir, T. W., et al., Biochem., 33:7701 (1994); McConnell, 30 S. J. and Hoess, R.H., JMB 250:460-470 (1995); Ku, J., et al., Proc. Natl. Acad. Sci., USA, 92:6552 (1995); Martin, F., et al., EMBO J.:5303-5309 (1994); Venturini, S., et al., Proteins and Peptides Lett., 1:70 (1994); Mutter, M., Ang. Chem. Int. Ed. Engl., 24:639 (1985); Cochran; A. G. 35 and Kim, P.S., Science, 271:1113-1116 (1996); O'Shea, E.

K., et al., Cell, 68:699 (1992); Oas, T. G., et al., Nature, 336:42 (1988).

Identification and production of the macromolecules of non-natural handedness which bind to the target 5 macromolecule can be carried out using any collection of macromolecules of natural handedness. The method described is applicable to all situations in which a biologically encoded library is used to isolate structures that interact with a chiral target (or chiral "bait") (Scott, J.K. and 10 Smith, G.P., Science 249:386 (1990); Devlin, J.J., et al., Ibid. 249:404 (1990); Cwirla, S.E., et al., Proc. Natl. Acad. Sci. USA 87:6378 (1990); Cull, M.G., et al., Proc. Natl. Acad. Sci. USA 89:1865 (1992); Mattheak, L.C., et al., Proc. Natl. Acad. Sci., USA 91:9022 (1994). As 15 described in the exemplification, a biologically encoded library, such as a phage display library (mirror image phage display) can be used in the methods of the present invention. Because ribonucleotide and deoxyribonucleotides also contain chiral centers which are recognized by nucleases, this approach equally applies to both RNA libraries and DNA libraries (Bock, L.C., et al., Nature, 355:564 (1992)). Therefore, the types of libraries for which this approach is useful include RNA (e.g, SELEX), DNA (e.g., DNA library) and peptide libraries (e.g., in vitro 25 transcription/translation based libraries, mono- and polyvalent phage libraries and 'peptide on plasmid' libraries). Use of a DNA library in the method of the present invention is described in Example 5. Examination of the vast amount of structural space represented in these libraries can yield new ligands for proteins of biological and medical importance. Phages which specifically interact with the Denantiomer and the L-enantiomer of a naturally-occurring (wildtype) macromolecule have been isolated, as described in the exemplification.

The selection process of the macromolecule of natural handedness in the library bound to the enantiomer of the target macromolecule can be performed in an achiral solvent (e.g., water) or chiral solvent. Furthermore, the 5 interaction between the naturally occurring macromolecule and the enantiomer is unlikely to require any additional chiral cofactors. An example of a selection process that can be used is described in the exemplification. Modification of the selection process can be performed 10 using methods known in the art.

In a particular embodiment, D amino acid peptides which are ligands for a naturally-occurring L amino acid peptide are identified by the claimed method. Such D amino acid peptides can be produced to correspond precisely to 15 the L amino acid peptide (except the constituent amino acids are D, not L enantiomers) or can be modified, such as by a substitution, deletion or modification of one or more constituent amino acids or addition of one or more D amino acid, to produce derivatives of the D amino acid peptides identified. 20

In another embodiment, L nucleic acid sequences which bind to D nucleic acid sequences or L amino acid peptides are identified by the claimed methods. The L nucleic acid sequences are produced to correspond to the D nucleic acid 25 (except that the constituent nucleotides are L nucleotides) or can be modified, such as by substitution, deletion or modification of one or more of the constituent L nucleotides or addition of one or more L nucleotides, to produce derivatives of the L nucleic acid sequence.

Methods of producing a derivative of a macromolecule of non-natural handedness which binds a target macromolecule of natural handedness using the methods described herein and modifying the macromolecule of nonnatural handedness to produce a derivative thereof is also 35 encompassed by the present invention. As used herein, the term "derivative" includes macromolecules of non-natural handedness identified, for example, by the methods described herein, which bind to the target macromolecule and have been modified in such a manner that they differ from the macromolecule of non-natural handedness by the addition, deletion, substitution or alteration of one or more components.

In one embodiment, the derivative is a modified Dpeptide. Derivatives of the D-peptides of the present

10 invention include, for example, D-peptides having modified
or altered (e.g., enhanced, decreased) affinity,
specificity, membrane permeability, hydrophobicity,
lipophilicity, oral bioavailability and/or biological halflife. Strategies for producing derivatives of D-peptides

15 include, for example, modifying the peptide backbone by Nmethylation (Ostresh, J.M., et al., Proc. Natl. Acad. Sci.,
USA, 91:11138-11142 (1994); Drug Discovery Technologies,
C.R. Clark, eds., John Wiley & Sons, 1990) and/or producing
peptide mimetics (Cho, C.Y., et al., Science, 261:1303

20 (1993); Moran E.J., et al., Biopolymers, 37:213-219
(1995)).

Alternatively, the sidechains of the D-peptides can be modified by introducing non-natural amino acids and amino acid analogues (Combs, A.P., et al., J. Am. Chem. Soc., 118: 287-288 (1996); Rivier, J.E., et al., Proc. Natl. Acad. Soc., USA, 93:2031 (1995); Munroe, J.E., et al., Bioorganic & Medicinal Chem. Lett., 5:2897-2902 (1995)). In a particular embodiment, the D-peptide contains non-naturally occurring side chains. Examples of derivatives of D-amino acid peptides having modified side chains can be described using the following formula: NH2-CHR-COOH wherein R is a lower alkyl, which is defined herein as an alkyl group of about 1 to about 50 carbon atoms which can be straight or branched and can include one or more double or triple bonds (e.g., methyl, ethyl). The lower akyl can

20

be a substituted lower akyl such as a(n) -NH2, -OH, aryl (e.g., phenyl), heteroaryl (e.g., imidazole, indole), -COOH (e.g., arginine) group. In addition, the lower akyl can be an aryl (e.g., phenyl, naphthyl), a substituted aryl (e.g., 5 -OH, halogen), a heteroaryl or a substituted heteroaryl group.

In addition, the D-peptides of the present invention can be modified by generating cyclic or polycyclic derivatives wherein, for example, disulfide bonds are 10 replaced to optimize and restrict the conformation of the D-peptide (Katz, B. A., et al., J. Am. Chem. Soc., 117:8541 (1995); Ladner R. C., TIBTECH, 13:426-430 (1995)). For example, the D-amino acid peptide identified as described herein can be made cyclic as in the case with cyclosporin, via a peptide bond.

Further, derivatives of the D-peptides of the present invention can be coupled to or incorporated within carriers to improve membrane permeability and/or bioavailability using, for example, liposomes and/or lipid derivatives (Eichholtz, T., et al., J. Biol. Chem., 268(3):1982-1986 1982 (1993)) and/or peptidic and non-peptidic polymers (Drug Discovery Technologies, C.R. Clark, eds., John Wiley & Sons, 1990); Sheldon, K., et al., Proc. Natl. Acad. Sci., USA, 92:2056 (1995)).

In addition, the D-amino acid peptide identified by 25 the methods described herein can be derivatized by increasing the hydrophobicity of the D-amino acid peptide (e.g., alkylating, for example methylating the D-amino acid peptide backbone), particularly when the target is an 30 intracellular target. A benzene ring can be added to the side chains to increase hydrophobicity. Alternatively, the D-amino acid peptide can be incorporated into a drug delivery system such as a liposome to increase hydrophobicity of the peptide.

Further, the C or N terminus of the D-peptide can be modified with protective groups (See for example, Green, T.H. and Wuts, P.G.M., Protective Groups in Organic Synthesis, second edition, John Wiley & Sons, N.Y. (1991)). 5 For example, a lipophilic polymer can be bonded to the N or C terminal (e.g., polyethylene glycol ester at the C terminal). Alternatively, one or more side chains can be bonded to a lipophilic polymer such as a polyalkylene glycol (e.g., polyethylene glycol), or bonded to an ether 10 or ester linkage (e.g., forms an ester with the side chain carboxyl group of aspartic acid or glutamic acid). For example, the D-peptide can be modified by producing a polymer-D-peptide conjugate wherein the polymer is, for example, monomethoxpolyethylene glycol (PEG) and/or 15 polyoxyethylated glycerol (POG) (see Therapeutic Peptides and Proteins, Marshak, D. and Liu, D., eds., Cold Spring Harbor Laboratory (1989)).

In another embodiment, the derivative is an L-nucleic acid. Strategies described herein for modifying D-peptides can also be used to modify RNA and DNA of non-natural handedness identified as described herein in order to optimize their pharmacological properties (e.g., Green, L. S., et al., Chem. Biol., 2:683 (1995); Latham, J. A., et al., Nucleic Acids Res., 22:2817 (1994); Gold, L., et al., Ann. Rev. Biochem., 64:763 (1995)).

Synthetic and biologically encoded libraries have proven to be extremely useful for the identification of ligands and nucleic acid sequences for a large variety of macromolecules. Synthetic peptide libraries composed of (D)-amino acids have been favored over gene-based techniques such as phage display Scott, J.K. and Smith, G.P., Science 249:386 (1990); Devlin, J.J., et al., Ibid. 249:404 (1990); Cwirla, S.E., et al., Proc. Natl. Acad. Sci. USA 87:6378 (1990), 'peptide on plasmid' Cull, M.G., et al., Proc. Natl. Acad. Sci. USA 89:1865 (1992) and in vitro translation based systems Mattheak, L.C., et al.,

Proc. Natl. Acad. Sci., USA 91:9022 (1994) because the
resulting peptides are insensitive to proteolytic digestion
and fail to induce an efficient immune response.
Furthermore, peptides composed of (D)-amino acids can be
absorbed intestinally (Pappenheimer, J.R., et al., Proc.
Natl. Acad. Sci. USA 91:1942 (1994)). Cyclosporin A, an 11
residue cyclic peptide composed mainly of N-methylated and
(D)-amino acids, is a leading immunosuppressant and is
generally given orally (Ptachcins, R.J., et al., Clin.
Pharmacokinetics, 11:107 (1986)).

Although the screening of a (D)-amino acid library has recently led to the identification of a peptide with analgesic activity (Dooley, C.T., et al., Science 266:2019 (1994)), the proportion of sequence space that can be sampled in synthetic libraries is generally only a fraction of what can be attained through the use of biologically encoded systems. Because of this lower degeneracy, and more importantly, because of the lack of intermediate amplification steps, the use of synthetic libraries has not always been as successful as the use of phage display libraries in the identification of ligands.

Proteins composed of (D)-amino acids have a chiral specificity for substrates and inhibitors that is the exact opposite of that of the naturally occurring (L)-amino acid protein (Del Milton, R.C., et al., Science, 257:1445 (1992); Petsko, G.A., Ibid, 256:1403 (1992); Zawadzke, L.E. and Berg, J.M., J. Am. Chem. Soc., 114:4002 (1992)). Although in certain instances (D)-amino acid ligands can be obtained by either making the (D)-enantiomer of a natural ligand (Fisher, P.J., et al., Nature 368:651 (1994)), or by making the 'reverse' (D)-enantiomer (Jameson, B.A., et al., Nature 368:744 (1994); Guptasarma, TIBTECH, 14:42-43 (1996)), such methods have no general applicability (Brady, L. and Dodson, G., Nature 368:692 (1994); Chorev, M. and Goodman, M., TIBTECH, 14:43-44 (1996)); Guichard, G., et al., TIBTECH, 14:44-45 (1996)).

A more general method to obtain (D)-amino acid ligands could be the selection of peptides from a biologically encoded library using the (D)-enantiomer of a protein of interest. Because the (D)- and (L)-protein have a chiral specificity for substrates and inhibitors that is the exact opposite, the (D)-enantiomeric form of the phage-displayed peptides that interact with the (D)-protein will interact with the protein of the natural handedness.

The validity of this approach has been shown by the 10 isolation of phage that specifically interact with the Denantiomer of the SRC homology 3 domain (SH3 domain). As described in Example 1, to examine the possible use of the mirror image relationship between (L) - and (D) -proteins for the identification of (D)-amino acid ligands from phage libraries, a (D)-amino acid version of the SH3 domain of 15 the c-Src tyrosine kinase was synthesized. SH3 domains are 55-70 amino acid protein domains that are found in a variety of intracellular effector molecules (reviewed in Schlessinger, J., Curr. Opin. Genet. Dev., 4:25 (1994)). 20 Because c-SRC activity is essential for osteoclast-mediated bone resorption, interference with SRC function may be of value in the treatment of osteoporosis (Soriano, P. et al., Cell, 64:693 (1992); Lowe, C., Proc. Natl. Acad. Sci USA, 90:4485 (1993); Seymour, J.F., Science and Medicine, 2:48 (1995)). SH3 domains interact with sequence elements in their cellular targets that form type II poly-proline helices of 8 to 10 residues (Rickles, R.J., et al., EMBO J., 13:5598 (1994); Sparks, A.B., J. Biol. Chem., 269:23853 (1994); Cheadle, C., et al., Ibid, 269:24034 (1994); Yu, 30 H., et al., Cell, 76:933 (1994); Feng, S., et al., Science

266:1241 (1994); Lim, W.A., et al., Nature, 372:375 (1994). Mediating protein interactions in intracellular cell signalling proteins, and interference with the signalling of SH3 domain containing proteins would be desirable.

35 Although ligands or substrates for a variety of SH3 domains have been isolated from phage display libraries (Rickles,

R.J., et al., EMBO J., 13:5598 (1994); Sparks, A.B., J.

Biol. Chem., 269:23853 (1994); Cheadle, C., et al., Ibid,

39:24034 (1994)), the identification of such sequences from
a synthetic (L)-amino acid peptide library was possible

only with prior knowledge of the sequences of the preferred
ligands (Yu, H., et al., Cell, 76:933 (1994)). Thus, the
identification of (D)-amino acid ligands for SH3 domains
from synthetic libraries is unlikely to be successful, in
the absence of prior sequence or structure information
about potential ligands.

The L- and D-enantiomers of the chicken c-SRC domain were prepared by bacterial expression and chemical synthesis, respectively. The biotinylated, synthetic, 60-amino acid D-SH3 domain was refolded and purified by affinity chromatography, with a D-amino acid version of a known peptide ligand for the SH3 domain (Yu, H., et al., Cell, 76:933 (1994)). As expected, bacterially expressed L-SH3 was retained on an affinity column with the L-enantiomer of this peptide, but not with the D-enantiomer, which indicates that the interaction of the SH3 domain with its substrates is stereospecific.

A phage library was constructed in which random, 10residue peptide sequences were expressed at the NH₂terminus of the pIII protein of the bacteriophage fd

25 (Scott, J.K. and Smith, G.P., Science, 249:386 (1990).
Because many natural bioactive peptides, such as the
immunosuppressant cyclosporin and the tumor promoter
microcystin, are cyclic, the library was designed to
include a large number of sequences that have a propensity

30 for disulfide bond formation (Smith, G.P. and Scott, J.K.,
Methods Enzymol., 217:228 (1993)). When the L-SH3 domain
was used to screen this phage display library for
interacting peptide sequences, disulfide-free polyprolinetype sequences that have been identified by others were

35 isolated (Yu, H., et al., Cell, 76:933 (1994); Rickles,
R.J. et al., EMBO J., 13:5598 (1994); Sparks, A.B., et al.,

J. Biol. Chem., 269:23853 (1994); Cheadle, C., et al., ibid., p. 24034)).

When the same phage display was screened with the D-SH3 domain, a series of peptide sequences that showed no 5 obvious sequence similarity to the L-SH3-binding sequences was isolated and grouped in three classes (Table 1). peptides all interact with the substrate binding site of the SH3 domain as they were eluted with the (D) -YGGRELPPLPRF peptide (SEQ ID NO: 2). These phage-displayed 10 peptides that bind to the D-SH3 domain are characterized by a combination of conserved leucine and glycine residues and a conserved arginine or lysine residue. In contrast to the L-peptide ligands for the L-SH3 domain (Yu, H., et al., Cell, 76:933 (1994); Rickles, R.J. et al., EMBO J., 13:5598 15 (1994; Sparks, A.B., et al., J. Biol. Chem., 269:23853 (1994); Cheadle, C., et al., ibid., p. 24034)), the positively charged residues in the ligands for the D-SH3 domain are located in the middle of a stretch of conserved residues, which suggests that the mode of ligand binding is 20 different in the two forms. Furthermore, all ligands for the D-SH3 domain contain a pair of cysteine residues, a property that is not observed for the L-peptides that interact with the L-SH3 domain (Yu, H., et al., Cell, 76:933 (1994); Rickles, R.J. et al., EMBO J., 13:5598 (1994; Sparks, A.B., et al., J. Biol. Chem., 269:23853 25 (1994); Cheadle, C., et al., ibid., p. 24034)). The disulfide bond may increase the affinity of these peptides for the D-SH3 domain by reducing the number of possible conformers.

Confirmation that the D-amino acid enantiomers of the peptides expressed by these phage particles interact with the all-L-amino acid SH3 domain is carried out using standard binding and detection methods. As described in Example 2, a D-peptide denoted Pep-D1, which is the mirror image of one of the phage-displayed peptides that bind to the D-SH3 domain, was synthesized and its interaction with

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the bacterially expressed L-SH3 domain examined. indirect binding assay was used to verify that the (D)-SH3 domain binds to the substrate binding site of the (L)-SH3 domain.

As described in Example 3, heteronuclear magnetic resonance (NMR) experiments were performed on the 15Nlabeled SH3 domain in the absence and presence of Pep-D1 to determine the binding site of this D-peptide in the SH3 domain. Residues in the SH3 domain that interact with Pep-10 D1 were identified through changes in amide 1H or 15N chemical shifts upon the addition of the D-peptide ligand.

In all cases the ligands that are isolated through this procedure are significantly less or not susceptible to the mechanisms that impair the activity of their biological 15 counterparts. For example, the ligands isolated through the method described herein are significantly less or not susceptible to RNase and DNase activity for nucleotidebased ligands (Ashley, G.W., J. Am. Chem. Soc. 114:9731 (1992); Urata, H., et al., J. Am. Chem. Soc., 113:8174 20 (1991) and proteolysis and activation of an immune response for peptide-based ligands (Gill, T.J., et al., Nature 197:746 (1963); Mauer, P.H., J. Exp. Med., 121:339 (1965); Borek, F., et al., Biochem., J., 96:577 (1965); Janeway, C.A. and Sela, M., Immunology 13:29 (1967); Ditzis, H.M., 25 et al., Proteins, 16:306 (1993)).

The approach described here to isolate ligands that are not of the natural handedness is unique for the isolation of oligonucleotide-based ligands, as all the approaches used thus far lead only to the isolation of ligands of the natural handedness. Consequently, such 'conventional' ligands are susceptible to degradation by natural occurring enzymes. In contrast, the synthesis and screening of synthetic peptide libraries composed of Damino acids is technically feasible. However, both because 35 of the high number of compounds that can be screened in biologically encoded library systems (several orders of

magnitude higher than can be achieved for synthetic peptide libraries) and the beneficial effects of the intermediate amplification steps used, such systems when combined with the technology described here yield superior results.

Specifically, synthetic peptide based strategies have 5 an upper limit in degeneracy that is determined either by peptide solubility limits and detection limits for the assay used (for synthetic combinatorial libraries and other solution-based peptide libraries), or for solid phase based 10 libraries, by volume considerations. Secondly, the intermediate amplification steps that are used in biologically encoded library systems allow the identification of ligands in situations where background binding is high (in systems that do not employ 15 amplification steps a specific ligand will only be identified if it constitutes an easily detectable part of the total pool of recovered molecules after a single round of screening). Thirdly, phage display and other biologically encoded library systems allow for the maturation of affinities through mutation of the encoding 20 DNA through processes such as error-prone PCR. Finally, phage libraries can accommodate inserts of significant length as compared to synthetic peptide based libraries. This not only allows the possible isolation of ligands of a 25 different size class, but significantly increases the complexity of short ligands that are contained (as sliding windows) within these inserts.

As described in Example 4, identification of D-amino acid peptide ligands which interact with a specific target as described herein can be used to provide guidelines for the design of biased (D)-amino acid peptide and peptidebased libraries. The libraries can subsequently be used to isolate novel ligands.

The invention is further illustrated in the following examples, which are not intended to be limiting in any way.

Example 1 Identification of Phage Which Specifically Interact With D-Amino Acid Peptides

Preparation of the L-SH3 domain

The residue numbering system is that of the full-5 length chicken c-SRC protein. Residues 81 to 140 of chicken c-SRC were cloned in other Hind III-Bam HI sites of the plasmid pMMHb (Staley, J.P. and Kim, P.S., Protein Science, 3:1822 (1994)). In this plasmid, proteins are expressed as a fusion with a modified form of the TrpLE 10 leader sequence in which the methionine residues have been replaced with leucine and the cysteine residues have been replaced with alanine (Staley, J.P. and Kim, P.S., Protein Science, 3:1822 (1994)), and a stretch of nine histidine residues has been inserted into the COOH terminal region of 15 the leader sequence. Expression of the fusion protein encoded by the plasmid pMMHb-SRC SH3 was induced at an absorbance of 0.6 at 600 nm by the addition of 0.4 mM $isopropyl-\beta-D-thiogalactopyranoside$ (IPTG) (Research Organics) to Escherichia-coli BL21-(DE3) pLys S cells 20 (Strategene). After induction for 2 hours, cells were centrifuged and inclusion bodies were isolated. Recombinant protein was purified by resuspension of inclusion bodies in 6 M guanidine-HCl and, 0.2 M tris, pH 8.7 (buffer A) and chromatography on a Ni^{2+} column (Ni^{-2+} 25 NTA-agarose; Qiagen). After elution, dialysis against water, and lyophilization, the fusion protein was dissolved in 70% formic acid and cleaved with CNBr (Stanley, J.P. and Kim P.S., Protein Science, 3:1822 (1994)). Dialyzed and lyophilized material was subsequently taken up in buffer A, and purified by chromatography on a Ni2+ column (after cleavage, the isolated SH3 domain flows through the column, whereas uncleaved fusion protein and the cleaved TrpLE leader sequence are retained). After dialysis (against PBS buffers of decreasing ionic strength, and finally against 35 water) and lyophilization, the purity and identity of the SH3 domain were confirmed by high-performance liquid

chromatography (HPLC) analysis at neutral pH and by laser desorption mass spectrometry (expected, 6686 daltons; observed, 6683 daltons).

Synthesis of the all D-Src SH3 domain

5 The all D amino acid SH3 domain, sequence GGVTTFVALYDYESRTETDLSFKKGERLQIVNNTEGDWWLAHSLTTGQTGYIPSNYVAP S-COOH-terminus (SEQ ID No: 1), residues 81-140 of chicken c-SRC, was synthesized on HMP resin (ABI/Perkin Elmer) with an ABI 431A peptide synthesizer and ABI fastmoc cycles 10 (Fmoc chemistry with HBTU activation and capping with acetic anhydride). Protected D-amino acids were obtained from Bachem California, Bachem Bioscience, Advanced Chemtech, and Novabiochem. For D-Ile and D-Thr, the side chain enantiomers were used, in which the chirality of the 15 side chain is also inverted relative to naturally occurring L-Thr and L-Ile. After completion of the synthesis, the NH2-terminus of the peptide was modified with NHS-LC-biotin II (Pierce). After cleavage, the peptide was lyophilized, dissolved in 6 M guanidine HCl, pH 6.0, and dialyzed 20 against 100 mM NaHPO4 and 100 mM NaCl, pH 6.0, with the use of dialysis tubing with a molecular cutoff of 3,500 daltons (D) (Spectra/Por). After dialysis the material was spun briefly to remove insoluble debris, and the supernatant was subsequently dialyzed against 5% acetic acid and 25 lyophilized. The peptide was dissolved at a concentration of 3.3 mg/ml in Tris-buffered saline (50 mM Tris, pH 7.5, and 150 mM NaCl) containing 1 mM biotin. biotinylated 60 amino acid (D)-SH3 domain length product was refolded and purified by affinity chromatography on an 30 all-(D) version of a known substrate for the Src SH3 domain (D)-YGGRELPPLPRF (SEQ ID NO:2), Yu, H., et al., Cell, 76:933 (1994)), that was biotinylated and immobilized on a streptavidin-agarose column (Pierce). This peptide is a derivative of an all (L)-peptide shown to bind to the all (L)-SH3 domain (Yu, H., et al., Cell, 76:933-945 (1994)),

with an NH2-terminal YGG added to facilitate concentration determination (H. Edelhoch, Biochemistry, 6:1948 (1967)). The L-peptide with the same NH2-terminal YGG served as a control ligand in the experiments. As expected,

5 bacterially expressed (L)-SH3 (residues 81-140 of chicken c-Src were expressed) can be retained on the (L)-enantiomer but not on the (D)-enantiomer of this peptide, indicating that the interaction of the SH3 domain with its substrates is stereospecific.

Chromatography fractions were analyzed by laser desorption mass spectrometry on a Voyager mass spectrometer (Perceptive Biosystems). Fractions containing material of the expected mass (expected, 7027 daltons; observed 7027 daltons to 7035 daltons) were pooled and dialyzed against 15 water for 72 hrs, lyophilized and taken up in water at a concentration of 107 µg/ml.

Production of phage library

The phage library was designed to provide expression of random peptides as NH2-terminal fusions with filamentous and the second of the seco 20 phage pIII protein. Typically, 3-5 copies were present per phage particle, to permit isolation of low/intermediate affinity ligands.

DNA encoding a 10-residue random insert with flanking serine or cysteine residues (S/C-X₁₀-S/C) (SEQ ID NO:20) 25 was prepared by PCR of an 85 residue oligonucleotide (Smith, G., "Cloning in Fuse vectors", Division of Biological Sciences, University of Missouri (Edition of February 10, 1992)) using biotinylated primers as described (Smith, G.P. and Scott, J.K., Methods Enzymol., 217:228 30 (1993)).

The insert design was: NH₂-A-D-G-A-S/C-X₁₀-S/C-G-A-G-A-PIII (SEQ ID NO:3).

85 Residue Oligonucleotide:

5'-C.TAT.TCT.CAC.TCG.GCC.GAC.GGG.GCT.TSC.(NNS)₁₀.TSC.GCC.GC
T.GGG.GCC.GAA.ACT.GTT.GAA-3' ((SEQ ID NO: 4)
In which S = C/G
N = A/T/C/G

5 (equimolar mixtures)

After purification of the PCR product and digestion with Bgl I, the end pieces were removed through the use of streptavidin-coated agarose beads (Pierce). The PCR product was subsequently ethanol precipitated and analyzed by electrophoresis on polyacrylamide gel.

The library was made by ligation of a random PCR product into Sfi I-cut Fuse 5 vector. After ligation the reaction mixture was extracted with phenol and chloroform, ethanol-precipitated and taken up in 10 mM Tris/1m mM EDTA 15 (pH 8.0). The ligation product was subsequently transferred into electrocompetent MC1061 cells (Biorad) and the Market and the using a Bio-Rad E. coli pulser and 0.1 cm cuvettes. After non-restrictive growth for 1 hr aliquots of transformed cells were plated on tetracycline-containing plates to the work placed 20 determine the efficiency of transformation, yielding an initial library of 3.6x108 transformants. transformation mixture was subsequently diluted to a volume of 400 ml LB and 20 $\mu \text{g/ml}$ tetracycline, and grown for an additional 14 hrs. A phage stock was prepared by two 25 successive polyethylene glycol (PEG) precipitations of the culture supernatant as described. The phage stock was finally resuspended in tris buffered saline/NaN, and titered by infection of K91-kan cells (A21), yielding a total of 2.8x1011 transforming units. The randomness of 30 the inserts was confirmed by sequencing of individual clones. 4x1010 transforming units were subsequently used to infect K91-kan cells to generate an amplified library. After 18 hrs phage were purified by three repetitive PEG precipitations yielding 10 ml of the amplified phage library (1.2-1012 transforming units/ml) in TBS/NaN3. The

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quality of the library was confirmed by selection of phage that expressed inserts that interact with 1) the lectin concanavalin A (Con A) (Oldenburg, K.R. et al., Proc. Natl. Acad. Sci. USA, 89:5393 (1992); Scott, J.K. et al., ibid, p. 5398); 2) two mouse monoclonal antibodies raised against the mouse MHC class I heavy chain; and 3) a bacterially expressed form of the Src SH3 domain, all screens giving the expected results.

Con A screen

10 After 3 rounds

AS W R Y N Y A F M R Y SA (SEQ ID No: 5)
(1)

AS M W M Y P Y P W G V SA (SEQ ID No: 6)
(9)

15 Screening of the phage library

The phage display was screened with the D-SH3 domain and a series of peptide sequences, which showed no obvious sequence similarity to the L-SH3-binding sequences, were isolated (see Table 1).

- Single wells of a flatbottom 96 well high binding E.I.A./R.I.A. plate (Costar) were coated overnight with 10 μg streptavidin (Pierce) in 100 μl 100 mM NaHCO3 at 4°C. After a single wash with water, wells were incubated with 100 μl (10.7 μg) of biotinylated (D)-SH3 for 1 hr. at 20°C, blocked for 2 hrs with 30 mg/ml dialyzed bovine serum albumin (BSA) in 100 mM NaHCO3, and again incubated with 100 μl (10.7 μg) of biotinylated (D)-SH3 for 1 hr. Unliganded streptavidin was blocked for 30 minutes by the addition of 8 μl 5 mM biotin in tris-buffered saline (TBS).
- Wells were subsequently washed 5 times with phosphate buffered saline (PBS) and 0.1% Tween-20, and incubated overnight with 50 μ l of the phage stock in TBS/NaN₃ and 50 μ l of TBS, 0.1% Tween-20, 1 mg/ml BSA and 0.05% NaN₃. Wells were subsequently washed by six additions of 200 μ l

of TBS, 0.1% Tween-20 and 1 mg/ml BSA with increasing incubation times in the later rounds of the selection procedure. (D)-SH3 bound phage particles were subsequently eluted by the addition of 100 μ l D-SH3 ligand peptide (715 5 μ M), sequence D-YGGRELPPLPRF-amide (SEQ ID No: 2), for 15 minutes at 4°C, at a final concentration of 700 to 1000 μM peptide. Acid elution of phages in this screen gives no detectable preferential binding to D-SH3 coated wells after four rounds of selection. The eluate was subsequently used 10 to infect K91-kan cells. Briefly 100 μ l eluate was mixed with 100 μ l K91 Terrific Broth cells (prepared as described) an incubated for 20 minutes at room temperature. The mixture was subsequently transferred into an Erlenmeyer flask containing 20 ml LB/0.2 μ g/ml tetracycline. After 1 15 hr incubation while shaking at 37°C, tetracycline was added to a final concentration of 20 $\mu g/ml$, appropriate dilutions were plated on tetracycline-containing plates ($20\mu g/ml$) to determine the titer of the eluate and the culture was incubated at 37°C for 12-16 hrs. Phage were isolated from 20 the supernatant by two PEG precipitations and the resulting phage stock was used for titering to determine the yield, and for the subsequent round of selection. In the fourth round of selection, phage were incubated in wells coated with or without the D-SH3 domain to determine the 25 specificity of the capture. The washing conditions and yields of the different rounds of selection were as follows:

	Round	Washing Conditions	<u>Yield</u>	
			D-SH3 coated	<u>Control</u>
30	1	6x, no incubation	1:3x10 ⁵	n.a.
	2	6x3 minutes at 4°C	1:1x10 ⁶	n.a.
	3	6x5 minutes at 4°C	1:2x104	n.a.
	4	6x10 minutes at 4°C	1:2x104	1:4x10 ⁵

(1994)).

From these data it is apparent that 1) in subsequent rounds phage that bind more tightly are selected since the yield increases although the washing conditions get significantly harsher and 2) this binding is specific for the D-SH3

- domain since elution is achieved by incubation with a substrate for this domain, and more importantly, since the recovery of phage particles in round 4 is 20-fold higher in the presence of the D-SH3 domain than in the absence (recovery is higher from the D-SH3 domain-coated plates).
- 10 A larger number of phage particles was retained in the presence of the D-SH3 domain than in its absence.

When the (D)-SH3 domain was used to screen this library a series of peptides were isolated that were grouped in three classes. These peptides all interact with

- the substrate binding site of the SH3 domain as they were eluted with the (D)-YGGRELPPLPRF peptide (SEQ ID NO:2). Surprisingly, all peptides contained a pair of cysteine residues, a property that is not observed for the polyproline peptides that interact with the (L)-SH3 domain.
- The disulfide bond may increase the affinity of these peptides for the (D)-SH3 domain by reducing the total number of possible conformers. Peptides in groups I and III contained at least a single arginine residue that may form a salt bridge with aspartic acid99 in the SH3 domain and which interacts with arginine residues in Src-binding poly proline peptides (Feng, S., et al., Science 266:1241)

Table 1. Sequences of phage-displayed peptides that interact with the (D)-Src SH3 domain.

Sequence	Number of Isolates		
	low stringency	high stringency	Туре
Group I			
CKRFVWRGQALC (SEQ ID No: 13)	10	14	
CSRASWRGLLFC (SEQ ID No: 14)		1	
Group II CWYLGYWPGQEC (SEQ ID No: 15)	12		
Group III			
CLSGLRLGLVPC (SEQ ID No: 16)	. 2		fdSRC-1
CLMGLRLGLLPC (SEQ ID No: 17)	4		fdSRC-1
CAYGFKLGLIKC (SEQ ID No: 18)	1*		fdSRC-3

This phage clone has an alanine to arginine substitution directly amino terminal to the insert region.

Conserved residues between different members of the groups I and III are indicated in bold, semiconserved residues are underlined. Note that for all members of group I and III the positioning of the conserved residues relative to the cysteine residues is preserved. Individual phage clones representing inserts of all 3 groups were analyzed after four to five rounds of selection for binding to the (D)-Src SH3 domain. All clones bind at least 150 fold better to wells coated with 0.5 µg streptavidin and 1.3 µg of the Src SH3 domain than to control wells. In both the low stringency and the high stringency screen, individual colonies were analyzed after 4 rounds of

selection. Both the high and low stringency screen were based on the same initial round of selection. Differences between the high stringency and low stringency screen include a 2 fold decrease in the concentration of (D)-Src SH3 used to coat the wells, a decrease in the incubation time of the phage on the plate (from 16 hrs. to 1 hr.), and an increase in the incubation time between the six washes of the plate (low stringency screen: round 2,3 and 4,3', 5' and 10' resp.; high stringency screen: rounds 2 to 4 all 10' incubations).

Phage display with the L-SH3 domain was also assessed. When the (L)-SH3 domain is used to screen a phage library (A12) for interacting sequences the poly-proline sequences that have been observed by others Rickles, R.J., et al., EMBO J., 13:5598 (1994); Sparks, A.B., J. Biol. Chem., 39:23853 (1994); Cheadle, C., et al., Ibid, 39:24034 (1994) are isolated. The results are as follows and showed that the display worked:

-33-

Bait

After 4 rounds

P P L V A P (SEQ ID No: 7) <u>P</u> E V L P I P (SEQ ID NO: 8) S R Α Α R L V P L R N (SEQ ID No: 9) 5 R M S P

GCN4 leucine zipper: short (33 residues), but low

probability

c-Src SH-3 domain: 60 residues, bind peptides

(type II polyPro)

10 GCN4 leucine zipper: synthesis straightforward, CD as

expected after 5 rounds no

difference in recovery +/- zipper

Test 59 individual clones, no

difference +/- zipper

15 Src SH3 domain: Src (L-) SH3 binds to substrate in

a stereo-specific manner

Src (L-) SH3 selects poly-Pro

sequences from library:

Sequence analysis of a small number of isolates after
four rounds of selection with the L-SH3 domain revealed the
following two peptide sequences: CLARSRLPAIPS (SEQ ID
NO: 10) (nine isolates) and SRMSPLVPLRNS (SEQ ID NO: 21)
(one isolate). The sequences of these peptides have
features consistent with those described for class I and
class II ligands of the SH3 domain ((Yu, H., et al., Cell,
76:933 (1994); Rickles, R.J. et al., EMBO J., 13:5598
(1994; Sparks, A.B., et al., J. Biol. Chem., 269:23853
(1994); Cheadle, C., et al., ibid., p. 24034)).

Analysis of single phage clones

To analyze the specificity of this interaction more rigorously, 6 individual colonies obtained after 4 rounds of selection were grown up in LB/20 μ g/ml tetracycline. 5 Phage particles were isolated from 1.3 ml supernatant by PEG precipitation and were resuspended in 500 5 μ l TBS giving an estimated concentration of 6.5x1010 transforming units/ml. 0.2 μ l aliquots (approximately 1.3x10 7 TU) were incubated in 50 μ l TBS/0.1% Tween 20/1 mg/ml BSA in wells 10 that had been coated as described above but with 0.5 rather than 10 μ g streptavidin and with or without 1.3 μ g D-SH3. After the incubation with phage non-bound phage were removed by 6(3 minute), washes with 150 μ l TBS/0.1% Tween20/1 mg/ml BSA. Bound phage particles were 15 subsequently eluted with 40 μ l glycinecine HCl pH2.2/1 mg/ml BSA for 10 minutes at 4°C. The eluate was subsequently brought to neutral pH and titered on K91 kan cells as described above.

	Clone	Recover (arbit:	rary units*	Ratio D-SH3/control
20		D-SH3 coated	<u>control</u>	
	1	1664	5	333
	2	1840	1	1840
	3	1316	0	>1316
	4	2348	0	>2348
25	5	1472	3	491
	6	1348	9	150

*Calculated from the number of colonies/plate at a given dilution.

Individual clones were analyzed after four and five rounds of selection. In subsequent rounds, the incubation time between washes was increased (times of 0, 3, 5, 10 and 10 min, respectively, for rounds 1 through 5). After four rounds of selection, 29 clones were sequenced, of which only 7 are within Group III of Table 1. To ensure that the selected phages were not binding to streptavidin or to a

composite surface formed by streptavidin formed by the streptavidin-D-SH3 complex, a fifth selection round was performed with neutravidin (Pierce) as a matrix. analysis of clones after this fifth round of selection 5 revealed only sequences of the fdSRC-2-type. Preliminary experiments suggest that the affinity of the corresponding peptide, Pep-D2, is similar to that of Pep-D1. Pep-D1 corresponds to the fdSRC-1 insert CLSGLRLGLVPC (SEQ ID No: 16) (Table 1), with the COOH-terminal alanine that is present in all flanking sequences (see Example 2). other phage isolates obtained after four rounds of selection expressed one of the following two sequences: CKRFVWRGQALC (SEQ ID No: 13) (10 isolates) and CWYLGYWPGQEC (SEQ ID No: 15) (12 isolates). The first of these sequences resembles the background sequences that are isolated with a variety of biotinylated ligates (Smith, G. P. and Scott, J.K., Methods in Enzymol., 217:228 (1993)) and is also similar to a sequence that was isolated previously with a monoclonal antibody against myohemerythrin, although it does not conform to the 20 recognition motif for this antibody (Smith, G. P. and Scott, J.K., Science, 249:386 (1990)). This sequence is therefore likely to bind to some component in the system other than the SH3 domain. Indeed, a D-amino acid version 25 of this sequence fails to bind to the L-SH3 domain, as judged by ELISA and NMR studies. The other sequence that was picked up after four rounds of selection shows limited similarity to the first sequence and has not been examined further.

30 Example 2 An All D Amino Acid Src SH3 Domain Binds to the L Src SH3 Domain

The (D)-amino acid peptide denoted Pep-D1, (D)RCLSGLRLGLVPCA (SEQ ID NO:11, a representative sequence of
group III sequences), which is the mirror image of one of
the phage-displayed peptides that binds to the D-Src SH3

domain, was synthesized and its interaction with the bacterially expressed (L)-SH3 domain was examined. Pep-D1 corresponds to the fdSRC-1 insert CLSGLRLGLVPC (SEQ ID No: 16) (Table 1), with the COOH-terminal alanine that is 5 present in all flanking sequences. The arginine immediately preceding the first cysteine residue was observed in the fdSRC-3 sequence (Table 1). The presence of arginine and lysine residues close to the NH2-terminus of secretory and transmembrane proteins negatively affects protein translocation (Boyd, D. and Beckwith, J., Cell, 62:1031 (1990)). In addition, a selection against arginine residues in the NH2-terminal part of phage pIII fusions has been observed (Cunningham, B.C. et al., EMBO J., 13:2508 The alanine to arginine mutation in this clone 15 may thus increase the affinity of the insert sequence for the D-SH3 domain, and could improve the solubility of the peptide; it was therefore included in the synthetic 2000 200 2000 peptide. For affinity measures, an NH2 terminal D-tyrosine was added to the peptide for concentration determination 20 (Edelhoch, H., Biochemistry, 6:1948 (1967). The peptides with and without the NH2 terminal tyrosine were airoxidized in 100 mM tris, pH 8.5, for 48 hours at a concentration of 1 mg/ml. Oxidized peptide was purified by reverse-phase HPLC with a C18 column and a wateracetonitrile gradient in 0.1% trifluoroacetic acid. identify of the products was confirmed by laser desorption mass spectrometry.

The reduced form of Pep-D1 shows no detectable binding activity in this assay (K_d>> 800 μM), which indicates that the formation of the disulfide is required for efficient binding. The affinity of Pep-D1 for the L-SH3 domain was determined by a competitive enzyme-linked immunosorbent assay (ELISA). Single wells of a 96-well plate were coated with 5 μg of the L-SH3 domain (Scott, J.K. and Smith, G.P., Science, 249:386 (1990); Smith, G.P. and Scott, J.K., Methods in Enzymol., 217:228 (1993)). Wells were blocked

with BSA, and phages expressing the L-SH3-binding insert CLARSRLPAIPS (SEQ ID NO: 10) were allowed to bind in 10 mM NaHPO4, pH 7.2, 15 mM NaCl, 1 mg/ml of BSA, 0.05% NaN3, and 0.1% Tween 20, in the presence of increasing amounts of 5 competitor peptide. Phage binding was quantified with a rabbit M13 antibody (Stratagene) and alkaline phosphataselabeled goat antibody to rabbit (Pierce), with a fresh solution of p-nitrophenol phosphate as substrate. Absorbance at 410 nm was determined with a Dynatech micotiter plate reader. Titration curves (means of triplicates) were obtained for the L-peptide ligand YGGRELPPLPRF amide (SEQ ID NO: 2) and the D-peptide ligand Pep-D1 YRCLSGLRLGLVPCA (SEQ ID NO: 22) in the presence and absence of 25 mM dithiothreitol. Relative values for $K_{\rm d}$ 15 were obtained as described (Minor, D.L., Jr. and Kim, P.S., Nature, 367:660 (1994)). The K_d of the L-peptide YGGRELPPLPRF-amide (SEQ ID NO: 2) was determined to be a second of the s 6.0 μ M by direct tryptophan fluorescence spectroscopy. A solution of the peptide was titrated into 1 μM SH3 solution 20 in 15 mM NaCl and 10 mM NaHPO, pH 7.2. Tryptophan fluorescence was induced by excitation at 295 nm (5 nm slit width), and emission was measured at 339 nm (10 nm slit width), with a Hitachi F-4500 fluorescence spectrometer. The dissociation constant was determined by Scatchard 25 analysis.

Although the syntheses of the (D)-enantiomeric form of both rubredoxin (45 amino acids) and human immunodeficiency virus (HIV) protease (99 amino acids) have been described Del Milton, R.C., et al., Science, 256:1445 (1992); Petsko, 30 G.A., Ibid, 256:1403 (1992); Zawadzke, L.E. and Berg, J.M., J. Am. Chem. Soc., 114:4002 (1992); Zawadzke, L.E. and Berg, J.M., Proteins, 16:301 (1993)), for most proteins the synthesis of the full (D)-enantiomeric form will not be feasible because of size limitations on the likelihood of successful chemical synthesis. However, both intracellular and extracellular proteins are often composed of autonomous

domains of 100 amino acids or less (reviewed in Bork, P. and Bairoch, A., Trends Biochem. Sci., 20, poster (1995) for extracellular proteins; Doolittle, R.F., and Bork, P., Sci. Am., 269:50 (1993); Efimov, A.V., FEBS Lett., 355:213 (1994); Cohen, G.B., et al., Cell, 80:237 (1995)). This size range is within reach of current solid-phase peptide synthesis technology, and recent advances in chemical ligation strategies for unprotected protein fragments hold promise for the synthesis of even larger protein domains. Thus, the isolation of ligands for proteins of interest (e.g., multidomain proteins) may be achieved through the synthesis and screening of one of its constituent domains, as described here for the SH3 domain.

Example 3 Determination of the Binding Site of the D-Peptide in the SH3 Domain

Heteronuclear magnetic resonance (NMR) experiments
were performed on the ¹⁵N-labeled SH3 domain in the absence
and presence of Pep-Dl to determine the binding site of
this D-peptide in the SH3 domain. Residues in the SH3

20 domain that interact with Pep-Dl were identified through
changes in amide ¹H or ¹⁵N chemical shifts upon the addition
of the D-peptide ligand.

The ligand-binding site of the SH3 domain for its natural, L-amino acid ligands consists of three pockets

25 that together form a relatively shallow groove on one side of the molecule (Feng, S., et al., Science, 266:1241 (1994); Yu, H., et al., Science, 258:1665 (1992)). Pocket A, which is formed by the side chains of aspartic acid and tryptophan accommodates the conserved arginine residue, whereas pockets B and C form a hydrophobic surface that accommodates the aliphatic and proline residues in SH3 ligands (Feng, S. et al., Science, 266:1241 (1994); Yu, H. et al., Science, 258:1665 (1992)).

Uniformly (\geq 395%) ¹⁵N-labeled SH3 domains were obtained 35 by growing *E. coli* harboring the plasmid pMMHb-SRC SH3 in

77 (45)

150

M9 medium supplemented with (15NH4)2SO4 (99.7% 15N; Isotec, Miamisburg, Ohio). Upon reaching an absorbance of 0.6 at 600 nm, cells were induced for 4 hours with 0.4 mM IPTG. The protein was purified as described for the unlabeled 5 material. Spectra were collected on a Bruker AMX 500 MHz NMR spectrometer. Resonance assignments were made by standard methods (Wüthrich, K., NMR of Proteins and Nucleic Acids (Wiley, New York, 1986); McIntosh, L.P. et al., Biochemistry, 29:6341 (1990)) and were consistent with the assignments for SH3 (Yu, H. et al., FEBS LETT., 324:87 (1993)). The peptide Pep-D1 was added to solution containing the 15N-labeled SH3 domain to a ratio of 1.5:1 (peptide:protein) in 10 mM phosphate, pH 6.0, at 298 K; heteronuclear single quantum coherence (HSQC) spectra (Bodenhausen, G. and Rubin, D.J., Chem. Phys. Lett., 69:185 15 (1980) of the uncomplexed and complexed forms were compared. There were no resonances with chemical shift differences >0.04 p.p.m. i the 1H dimension, or >0.17 p.p.m. in the 15N dimension. However, a number of resonances were reduced in intensity or completely absent in HSQC spectra of the complex. Residues that had the intensity of their HSQC resonances reduced significantly upon Pep-D1 binding, as compared to the ligand-free spectra, were identified as follows: for individual peaks, 25 the ratio of peak intensities in the absence and presence of peptide was determined and converted to a log scale. The resulting distribution around the median is markedly skewed toward the left. A window that included >90% of the residues with ratios that were higher than the median was 30 applied to residues with chemical shifts below the median. Only residues with a ratio lower than the median and that were not contained within this window were considered to have undergone significant perturbation (according to these criteria, only residues with a ratio that was reduced to less than 0.65 of that of the median were considered to have undergone significant perturbation). These residues

1.

include residues 94, 97, 112, 115, 117, 119, 120, 131, 132 and 135, the indole resonance of tryptophan119, and the side chain amides of asparagine¹¹³, and asparagine¹³⁵. The resonances of 95, 96, 98, 99, 100, 118 and 134 and the 5 indole resonance of tryptophan were absent in the presence of ligand. Control experiments, using the Lpeptide YGGRELPPLPRF amide (SEQ ID NO: 2) resulted in 17 resonances that were shifted by ≥30.1 p.p.m. in the ¹H dimension or \ge 30.5 p.p.m. in the 15N dimension (residues 87, 10 89, 90, 92, 96, 98, 99, 100, 109, 111, 114, 116, 119, 121, 131, 132 and 135, the indole resonance of tryptophan¹¹⁹, and the side chain amides of asparagine¹¹³ and asparagine¹³⁵). Five resonances (95, 97, 117, 118 and 134) were absent in HSQC spectra of the complex. To validate the approach 15 chosen to identify residues that interact with Pep-D1, this approach was applied to the spectra obtained with the peptide L-YGGRELPPLPRF amide (SEQ ID NO: 2). With this approach, no new residues were identified that interacted with this peptide. The effect of peptide binding on the 20 chemical shift of proline 133, which forms part of pocket B, could not be observed in this type of experiment.

> The binding of Pep-D1 results in the perturbation of the chemical shifts of the residues that form pocket A, as well as a patch of adjacent residues (Fig. 2C). Most of 25 these residues also undergo changes in their chemical shifts upon binding of the L-peptide (Fig. 2B). Pocket A is likely to interact with the conserved arginine or lysine residues in the D-peptides in a manner that is analogous to the recognition of arginine residues in L-amino acid ligands. The interaction of this site with both the L- and D-amino acid ligands explains the competition observed for the binding of these two ligands.

Pep-D1 appears to occupy only part of the binding site that is contacted by the polyproline-type ligands for the SH3 domain (Fig. 3). Residues that form part of pocket B and pocket C (tyrosine90 and tyrosine92), or that are

adjacent to this pocket (valine⁸⁷ and leucine⁸⁹), are not perturbed upon binding of Pep-D1 (Figs. 2 and 3).

Mutational analysis suggests that, for L-amino acid ligands, interactions at these sites are required for high-affinity binding (Feng, S. et al., Science, 266:1241 (1994). D-peptide inhibitors of higher affinity could therefore potentially be obtained by the design or selection of analogs of Pep-D1 or Pep-D2 (Table 1) that extend further along the groove, into pocket C of the SH3 domain.

Example 4 Use of the D-amino acid peptide ligands to design highly-enriched libraries

Random synthetic libraries do not cover enough conformational space to always allow for the isolation of 15 high affinity ligands for a given target. A more promising strategy is the use of libraries that are biased towards structural elements known to interact with a target of Specifically, the D-amino acid peptide sequences that interact with a given target that are isolated using 20 the strategy described here are used as guidelines for the design of biased (D)-amino acid peptide and peptide-based libraries. These biased libraries may subsequently be used for the isolation of novel ligands. For example, the three classes of (D)-amino acid peptide ligands for the SH3 domain (Example 2) are useful to design (D)-amino acid peptide-based libraries highly enriched for SH3-binding peptides or peptidomimetics. Such libraries are useful to identify peptides which bind the SH3 domain and are particularly useful because they are biased toward (have an 30 enhanced content of) peptides known to bind the SH3 domain. All SH3 domains for which the interaction with (L)-amino acid peptides has been examined bind to ligands with similar structural elements. Synthetic libraries based on the structure of (D)-amino acid ligands for the SH3 domain 35 are also enriched in ligands for other SH3 domains. The

biased libraries based on the structure of the (D)-amino acid peptide ligands for the SH3 domain are useful for the isolation of ligands for a variety of SH3 domains.

A biased library is constructed, for example, as

follows, based on the amino acid sequences of the 3 classes
of peptides described in Example 2, which have been shown
to bind the (D)-SH3 domain. A chemical peptide library of
D-amino acids is prepared in which about 80% of the D-amino
acid peptides of the library have the conserved amino acid
residues and about 20% of the D-amino acid peptides of the
library do not have the conserved amino acid residues.

Thus, the library, which is heavily biased for peptides having the general structure of peptides known to bind (D)-SH3 domain (i.e., 80%), can be used to isolate other D-amino acid peptides which have the conserved structure and bind to other SH3 domains (i.e., human). In addition, D-amino acid peptides in which the conserved amino acid residue has been altered (i.e., from the 20% of the D-amino acid peptides) and which binds to the SH3 domain with equivalent or greater affinity, can be isolated.

Example 5 Mirror Image Selection of an Enantiomeric DNA Inhibitor of Vasopressin

Vasopressin was prepared using conventional solid 25 phase peptide synthesis.

In the first step toward generating an antagonist of the peptide hormone L-vasopressin, an in vitro selection was used to isolate single-stranded DNAs (ssDNAs) that bind synthetic D-vasopressin (DVP). A starting pool of 10¹⁶

30 different 96-mers was synthesized on a DNA synthesizer. Each pool molecule contained a central region with 60 random-sequence positions that was flanked by two 18-nt defined regions. Molecules within this starting pool that bind DVP were enriched by affinity chromatography. The

affinity resin was prepared by coupling biotinylated DVP to streptavidin agarose. The ssDNA pool was radiolabeled, denatured, renatured in a physiological buffer and passed over this resin. After extensively washing the column, 5 molecules that bind DVP were eluted from the column with a molar excess of DVP. Eluted pool was amplified by PCR using a negative-strand "primer-terminator", that is, an oligonucleotide bearing a central segment of non-nucleotide material that blocks further extension of the positive 10 strand (Williams, K.P. and Bartel, D.P., Nucleic Acids Res., 23:4220-4221 (1995)). Such PCR results in a substantial size difference between the two product strands, facilitating gel-purification of the positivestrand ssDNA pool. This combination of affinity 15 chromatography and PCR constituted one cycle of selection amplification.

After thirteen iterations of the selectionamplification cycle the pool was cloned. Analysis of 56
clones revealed only two different sequences, 96.2 and
20 96.4. Positive ssDNA of each sequence binds DVP and both
have potential to form very similar secondary structures.
Affinity chromatography studies of a set of deletion
mutants has identified one, 69.1, that appears to bind DVP
better than does the parent sequence. This aptamer is not
eluted from the DVP resin by L-vasopressin.

96.2 TCTAACGTGAATGATAGAcggcgaatccccaatgcgaagcagtggttttgca GTCGAGTTGCTGTGCCGATGAgcgTTAACTTATTCGACCAAA (SEQ ID NO:23)

30 96.4

TCTAACGTGAATGATAGAcgttacgtgtctacactatGTCGAGTTGCTGT

GTGCCGATGAacgtgggattagagcgtgTTAACTTATTCGACCAAA (SEQ ID
NO:24)

69.1
TCTAACGTGAATGATAGACGttacgtgtctacactatGTCGAGTTGCTGT
GTGCCGATGAacgtgggat (SEQ ID NO:25)

D-vasopressin aptamers. Sequence blocks shared by the two original aptamers are in capital letters. The defined-sequence segments that flanked the random-sequence region are indicated in bold. A segment derived from random-sequence positions that is shared by the two sequences of the final pool is in outline. The 69.1 aptamer is a deletion derivative of 96.4.

The information required to complete the selectionreflection procedures is described above. The synthesis of
the 69.1 sequence using enantio-deoxyribose
phosphoramidites will yield an aptamer that binds natural
vasopressin but is not susceptible to nuclease degradation.
However, before synthesizing such an aptamer, a second
selection experiment can be performed, starting with a
degenerate set of sequences based on the 69.1 sequence.
This allows for isolation of sequence variants that bind
DVP with higher affinity. Data from this second selection
will also provide a reliable secondary structure model that
will guide further deletion experiments, yielding a final
DVP aptamer of smaller size and greater activity. This
nuclease-proof aptamer will be tested in vitro as a ligand
and in vivo as an antagonist of vasopressin.

Equivalents

Those skilled in the art will recognize, or be able to ascertain using no more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following claims.

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 - (A) MEDIUM TYPE: Floppy disk

 - (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
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 - (B) FILING DATE:
 - (C) CLASSIFICATION:
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 - (A) APPLICATION NUMBER: US 08/627,497
 - (B) FILING DATE: 28-MAR-1996
 - (C) CLASSIFICATION:

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 - (A) APPLICATION NUMBER: US 60/001,067
 - (B) FILING DATE: 11-JUN-1995
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/482,309
 - (B) FILING DATE: 07-JUN-1995
 - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/433,572
 - (B) FILING DATE: 03-MAY-1995
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
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 - (B) REGISTRATION NUMBER: 32,227
 - (C) REFERENCE/DOCKET NUMBER: WHI95-04A3
 - (ix) TELECOMMUNICATION INFORMATION:
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 - (B) TELEFAX: 617-861-9540
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 60 amino acids

 - (B) TYPE: amino acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:
 - Gly Gly Val Thr Thr Phe Val Ala Leu Tyr Asp Tyr Glu Ser Arg Thr
 - Glu Thr Asp Leu Ser Phe Lys Lys Gly Glu Arg Leu Gln Ile Val Asn
 - Asn Thr Glu Gly Asp Trp Trp Leu Ala His Ser Leu Thr Thr Gly Gln
 - Thr Gly Tyr Ile Pro Ser Asn Tyr Val Ala Pro Ser 55 50
- (2) INFORMATION FOR SEQ ID NO:2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 (B) TYPE: amino acid

 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Tyr Gly Gly Arg Glu Leu Pro Pro Leu Pro Arg Phe

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (ix) FEATURE:
 - (A) NAME/KEY: Modified-site
 - (B) LOCATION: 5
 - (D) OTHER INFORMATION: /product= "OTHER"
 - /note= "The Ser at this location can be either Ser or Cys."
 - (ix) FEATURE:
 - (A) NAME/KEY: Modified-site
 - (B) LOCATION: 16
 - (D) OTHER INFORMATION: /product= "OTHER"

/note= "Ser at this location can be either Ser or Cys."

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Ala Asp Gly Ala Ser Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Ser

Gly Ala Gly Ala 20

- (2) INFORMATION FOR SEQ ID NO:4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 85 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (other)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

CTATTCTCAC TCGGCCGACG GGGCTTSCNN SNNSNNSNNS NNSNNSNNSN NSNNSNNSTS

60 85

CGCCGCTGGG GCCGAAACTG TTGAA

- (2) INFORMATION FOR SEQ ID NO:5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids
 (B) TYPE: amino acid

 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Ala Ser Trp Arg Tyr Asn Tyr Ala Phe Met Arg Tyr Ser Ala

- (2) INFORMATION FOR SEQ ID NO:6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Ala Ser Met Trp Met Tyr Pro Tyr Pro Trp Gly Val Ser Ala 10

- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Leu Pro Glu Val Pro Pro Leu Val Ala Pro 1

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Leu Ala Arg Ser Arg Leu Pro Ala Ile Pro

- (2) INFORMATION FOR SEQ ID NO:9:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 10 amino acids

 - (B) TYPE: amino acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

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- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Arg Met Ser Pro Leu Val Pro Leu Arg Asn

- (2) INFORMATION FOR SEQ ID NO:10:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Cys Leu Ala Arg Ser Arg Leu Pro Ala Ile Pro Ser

- (2) INFORMATION FOR SEQ ID NO:11:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids

 - (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Arg Cys Leu Ser Gly Leu Arg Leu Gly Leu Val Pro Cys Ala 5

- (2) INFORMATION FOR SEQ ID NO:12:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 14 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Ala Cys Lys Arg Phe Val Trp Arg Gly Gln Ala Leu Cys Ala

- (2) INFORMATION FOR SEQ ID NO:13:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single

- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: peptide
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Cys Lys Arg Phe Val Trp Arg Gly Gln Ala Leu Cys

- (2) INFORMATION FOR SEQ ID NO:14:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Cys Ser Arg Ala Ser Trp Arg Gly Leu Leu Phe Cys

- (2) INFORMATION FOR SEQ ID NO:15:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Cys Trp Tyr Leu Gly Tyr Trp Pro Gly Gln Glu Cys

- (2) INFORMATION FOR SEQ ID NO:16:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids

 - (B) TYPE: amino acid
 (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

Cys Leu Ser Gly Leu Arg Leu Gly Leu Val Pro Cys

- (2) INFORMATION FOR SEQ ID NO:17:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

Cys Leu Met Gly Leu Arg Leu Gly Leu Leu Pro Cys

- (2) INFORMATION FOR SEQ ID NO:18:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Cys Ala Tyr Gly Phe Lys Leu Gly Leu Ile Lys Cys

- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 11 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Cys Lys Arg Phe Trp Arg Gly Gln Ala Leu Cys

- (2) INFORMATION FOR SEQ ID NO:20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (ix) FEATURE:
 - (A) NAME/KEY: Modified-site
 - (B) LOCATION: 1
 - (D) OTHER INFORMATION: /product= "OTHER"
 - /note= "The amino acid at this location can also be Cys."

- (ix) FEATURE:
 - (A) NAME/KEY: Modified-site
 - (B) LOCATION: 12
 - (D) OTHER INFORMATION: /product= "OTHER"

/note= "The amino acid at this location can also be Cys."

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Ser Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Ser

- (2) INFORMATION FOR SEQ ID NO:21:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids

 - (B) TYPE: amino acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Ser Arg Met Ser Pro Leu Val Pro Leu Arg Asn Ser

- (2) INFORMATION FOR SEQ ID NO:22:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 15 amino acids

 - (B) TYPE: amino acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

Tyr Arg Cys Leu Ser Gly Leu Arg Leu Gly Leu Val Pro Cys Ala

- (2) INFORMATION FOR SEQ ID NO:23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 96 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (other)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

TCTAACGTGA ATGATAGACG GCGAATCCCC AATGCGAAGC AGTGGTTTTG CAGTCGAGTT GCTGTGTGCC GATGAGCGTT AACTTATTCG ACCAAA

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(2) INFORMATION FOR SEQ 12 NOTE:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 96 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (other)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:	
TCTAACGTGA ATGATAGACG TTACGTGTCT ACACTATGTC GAGTTGCTGT GTGCCGATGA	60
ACGTGGGATT AGAGCGTGTT AACTTATTCG ACCAAA	96
(2) INFORMATION FOR SEQ ID NO:25:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 69 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (other)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:	
TCTAACGTGA ATGATAGACG TTACGTGTCT ACACTATGTC GAGTTGCTGT GTGCCGATGA	60
ACGTGGGAT	69

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CLAIMS

What is claimed is:

- 1. A method of producing a macromolecule of non-natural handedness that binds to a target macromolecule of natural handedness, comprising the steps of:
 - a) providing an enantiomer of the target macromolecule or of a domain characteristic thereof;
 - b) providing a library of macromolecules of natural handedness;
 - c) contacting the library of b) with the enantiomer of a), under conditions appropriate for binding of a macromolecule of natural handedness in the library with the enantiomer of a), whereby the enantiomer of a) binds a macromolecule of natural handedness present in the library; and
 - d) producing the enantiomer of the macromolecule of natural handedness which is bound to the enantiomer of a),
- wherein the enantiomer of d) is a macromolecule of non-natural handedness which binds to the target macromolecule of natural handedness.
- 2. A method of producing a macromolecule of non-natural handedness that binds to a target macromolecule of natural handedness, comprising the steps of:
 - a) providing an enantiomer of the target macromolecule or of a domain characteristic thereof;
 - b) providing a library of macromolecules of natural handedness;
 - c) contacting the library of b) with the enantiomer of a), under conditions appropriate for binding of a macromolecule of natural handedness in the library with the enantiomer of a); whereby the

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- enantiomer of a) binds a macromolecule of natural handedness present in the library;
- d) identifying a macromolecule which is bound to the enantiomer of a);
- e) determining the sequence of the macromolecule of natural handedness identified in d); and
- e) producing a macromolecule of non-natural handedness which is the enantiomer of the macromolecule identified in d) or of a characteristic domain thereof,

wherein the enantiomer of e) is a macromolecule of non-natural handedness which binds to the target macromolecule of natural handedness.

- 3. The method of Claim 2 wherein the target macromolecule is a protein.
- 4. The method of Claim 3 wherein the protein is selected from the group consisting of: vasopressin, interleukin-8, thrombomodulin EGF-like domain, GPII_b-III_a cytoplasmic domain, Factor VIIa GLA domain, Factor IX EGF-like domain, HIV protease, NH₂-terminal SH3 domain GRB2, COOH-terminal SH3 domain GRB2, P120^{GAP} SH3 domain, vascular permeability factor and vascular endothelial growth factor.
- 5. The method of Claim 2 wherein the target macromolecule is an oligonucleotide.
 - 6. The method of Claim 5 wherein the oligonucleotide is selected from the group consisting of: HIV RRE, HIV Tar and BCR-AB1 fusion DNA sequences.
- 7. A method of producing a D amino acid peptide that binds to a target L macromolecule, comprising the steps of:

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- a) providing a D amino acid peptide of the target L macromolecule or of a domain characteristic thereof;
- b) providing a library of L amino acid peptides;
- c) contacting the library of b) with the D amino acid peptide of a), under conditions appropriate for binding of an L amino acid peptide in the library with the D amino acid peptide of a) whereby the peptide of a) binds an L amino acid peptide present in the library;
 - d) identifying an L amino acid peptide which is bound to the D amino acid peptide of a);
 - e) determining the sequence of the L amino acid peptide identified in d); and
- e) producing a D amino acid peptide of the L amino acid peptide identified in d) or of a characteristic domain thereof,

wherein the D amino acid peptide of e) binds to the target L macromolecule.

- 20 8. The method of Claim 7 wherein the peptide is selected from the group consisting of: vasopressin, interleukin-8, thrombomodulin EGF-like domain, GPII_b-III_a cytoplasmic domain, Factor VIIa GLA domain, Factor IX EGF-like domain, HIV protease, NH₂-terminal SH3 domain GRB2, COOH-terminal SH3 domain GRB2, P120^{GAP} SH3 domain, vascular permeability factor and vascular endothelial growth factor.
 - 9. A method of producing an L oligonucleotide that binds to a target L macromolecule, comprising the steps of:
- a) providing a D amino acid peptide of the target L macromolecule or of a domain characteristic thereof;
 - b) providing a library of D oligonucleotide;

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- contacting the library of b) with the D amino c) acid peptide of a), under conditions appropriate for binding of a D oligonucleotide in the library with the D amino acid peptide of a), whereby the peptide of a) binds a D oligonucleotide present in the library;
- identifying a D oligonucleotide which is bound to d) the D amino acid peptide of a);
- determining the sequence of the D oligonucleotide e) identified in d); and
- producing a L oligonucleotide of the D e) oligonucleotide identified in d) or of a characteristic domain thereof,

wherein the L oligonucleotide of e) binds to the target L macromolecule.

- The method of Claim 9 wherein the macromolecule is selected from the group consisting of: vasopressin, interleukin-8, Thrombomodulin EGF-like domain, GPII_b-III, cytoplasmic domain, Factor VIIa GLA domain, Factor IX EGF-like domain, HIV protease, NH2-terminal 20 SH3 domain GRB2, COOH-terminal SH3 domain GRB2, P120 GAP SH3 domain, vascular permeability factor and vascular endothelial growth factor.
- A method of identifying an L amino acid peptide which binds a D amino acid peptide of interest, comprising 25 the steps of:
 - providing a phage display library which comprises a) L amino acid peptides displayed on phage surfaces;
- contacting the phage display library of a) with 30 b) the D amino acid peptide of interest, under conditions appropriate for binding of L amino acid peptides displayed on phage surfaces with the D amino acid peptide of interest; and

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- c) identifying phages on the surfaces of which the D
 amino acid peptide of interest is bound to an L
 amino acid peptide displayed on the surface,
 thereby producing a D amino acid peptidedisplayed L amino acid peptide complex
 wherein the displayed L amino acid peptide is an L
 amino acid peptide which binds the D amino acid of
 interest.
- 12. The method of Claim 11 further comprising making a D

 amino acid peptide which corresponds to the L amino
 acid identified and further comprising the steps of:
 - d) determining the amino acid sequence of the L amino acid peptide displayed on the surface of the phage; and
- e) synthesizing the D amino acid peptide which corresponds to the amino acid sequence of the L amino acid peptide determined in d), thereby producing a D amino acid peptide which corresponds the L amino acid peptide displayed on the surface.
 - 13. The method of Claim 11 wherein the L amino acid peptides displayed on the phage surfaces bind D amino acid peptides of the src SH3 domain.
- 14. A synthetic amino acid peptide which binds the src SH3 domain.
 - 15. A synthetic D amino acid peptide corresponding to all or a portion of the src SH3 domain.
 - 16. A synthetic D amino acid peptide which binds a domain of an intracellular signaling protein.

- 17. A method of obtaining an L nucleic acid sequence which binds an L amino acid peptide of interest, comprising the steps of:
 - a) providing a collection of D nucleic acid sequences;
 - b) contacting the D nucleic acid sequences of a) with a D amino acid peptide of interest, under conditions appropriate for binding of the D nucleic acid sequences with the D amino acid peptide of interest;
 - c) isolate a D nucleic acid sequence which binds to the D amino acid peptide;
 - d) determine the nucleotide sequence of the D nucleic acid sequence of c); and
- e) prepare a nucleic acid sequence having the nucleotide sequence of d) using L nucleotides, wherein the nucleic acid sequence of e) is an L nucleic acid sequence which binds an L amino acid peptide.
- 20 18. A D amino acid peptide identified by the method of Claim 7.
 - 19. An L oligonucleotide identified by the method of Claim 9.
- 20. A macromolecule of non-natural handedness, produced by
 the method of Claim 1, that binds to a target
 macromolecule of natural handedness, wherein the
 target macromolecule is selected from the group
 consisting of: proteins, oligonucleotides and
 phospholipids.
- 30 21. A macromolecule of non-natural handedness of Claim 20, which binds to a target macromolecule and the target macromolecule is a protein selected from the group

consisting of: intracellular signaling proteins and domains thereof; chemokines; cytokines; enzymes; growth factors; growth factor receptors and domains thereof; transcription factors and domains thereof; and hormones.

A macromolecule of non-natural handedness of Claim 21 22. wherein the target macromolecule is selected from the group consisting of: SH3 domains; SH2 domains; PH domains; α -chemokine; β -chemokine; IL-1; TNF; lymphotoxin-α; IL-1β; IL-6; M-CSF; TGF; protein kinase 10 C; phospholipase C; phospholipase D; growth factors and growth factor receptors in the PDGF family, EGF family, FGF family, IGF family, HFG family, VEGF family, neurotrophin family, Eph family, Class I cytokine family, GH family, IL-3 family, IL-6 family, 15 IL-2 family, Class II cytokine family or TNF family; protein kinases; protein phosphatases; cyclins; Cdc proteins; Ets domain; bZIP; rel homology domain; STATs; NF-ATs; TCF; Fos; JAKs; human CD2; human CD58; human endothelin; heregulin- α ; human interleukin- 1β 20 converting enzyme; human macrophage inflammatory protein $1-\beta$; platet factor 4; human melanoma growth stimulating activity; GRO/melanoma growth stimulating activity; MHC molecules; bacterial muramidase; kringle domains; ras; ras-GAP; selections; Pleckstrin homology 25 domains; stromelysin; thrombin; tissue factor; calmodulin; CD4; collagenase; dihydrofolate reductase; fibronectin; fibronectin type III modules; G-protein subunits; vasopressin; Factor IX GLA domain; factor; interleukin-8; thrombomodulin EGF-like domain; GPIIb-30 III, cytoplasmic domain; Factor VIIa GLA domain; Factor IX EGF-like domain; human immunodeficiency virus (HIV) proteins; NH2-terminal SH3 domain GRB2; COOH-terminal SH3 domain GRB2; P120 GAP SH3 domain;

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vascular permeability factor and vascular endothelial growth factor.

- 23. A macromolecule of non-natural handedness of Claim 20, which binds to a target macromolecule and the target macromolecule is an oligonucleotide selected from the group consisting of: HIV RRE; HIV Tar; and BCR-AB1 fusion DNA sequences.
- A macromolecule of non-natural handedness of Claim 20 24. which binds to a target macromolecule and the target macromolecule is a phospholipid selected from the 10 group consisting of: phosphoinositide; phosphoinositidase C; phosphoinositide 3-kinase; phosphatidylinositol; phosphatidylinositol 3phosphate; phosphatidylinositol (4,5)bisphosphate; phosphatidylinositol (3,4,5)triphosphate; 15 . phosphatidylcholine; phosphatidylethanolamine; phosphatidic acid; inositol (1,4) bisphosphate; inositol (1,4,5) triphosphate; diacylglycerol; sphingosine; sphingosine phosphate; sphigosine phosphocholine and ceramide. 20
 - 25. A macromolecule of non-natural handedness, produced by the method of Claim 7, that binds to a target macromolecule of natural handedness, wherein the target macromolecule is selected from the group consisting of: proteins, oligonucleotides and phospholipids.
 - 26. A macromolecule of non-natural handedness of Claim 25, which binds to a target macromolecule and the target macromolecule is a protein selected from the group consisting of: intracellular signaling proteins and domains thereof; chemokines; cytokines; enzymes; growth factors; growth factor receptors and domains

thereof; transcription factors and domains thereof; and hormones.

A macromolecule of non-natural handedness of Claim 26 27. wherein the target macromolecule is selected from the group consisting of: SH3 domains; SH2 domains; PH 5 domains; α -chemokine; β -chemokine; IL-1; TNF; lymphotoxin- α ; IL-1 β ; IL-6; M-CSF; TGF; protein kinase C; phospholipase C; phospholipase D; growth factors and growth factor receptors in the PDGF family, EGF family, FGF family, IGF family, HFG family, VEGF 10 family, neurotrophin family, Eph family, Class I cytokine family, GH family, IL-3 family, IL-6 family, IL-2 family, Class II cytokine family or TNF family; protein kinases; protein phosphatases; cyclins; Cdc proteins; Ets domain; bZIP; rel homology domain; 15 STATs; NF-ATs; TCF; Fos; JAKs; human CD2; human CD58; human endothelin; heregulin- α ; human interleukin- 1β converting enzyme; human macrophage inflammatory protein $1-\beta$; platet factor 4; human melanoma growth stimulating activity; GRO/melanoma growth stimulating 20 activity; MHC molecules; bacterial muramidase; kringle domains; ras; ras-GAP; selectins; Pleckstrin homology domains; stromelysin; thrombin; tissue factor; calmodulin; CD4; collagenase; dihydrofolate reductase; fibronectin; fibronectin type III modules; G-protein 25 subunits; vasopressin; Factor IX GLA domain; factor; interleukin-8; thrombomodulin EGF-like domain; GPII_b-IIIa cytoplasmic domain Factor VIIa GLA domain; Factor IX EGF-like domain; human immunodeficiency virus (HIV) proteins; NH2-terminal SH3 domain GRB2; COOH-terminal 30 SH3 domain GRB2; P120 GAP SH3 domain; vascular permeability factor nadvascular endothelial growth factor.

- 28. A macromolecule of non-natural handedness of Claim 25, which binds to a target macromolecule and the target macromolecule is an oligonucleotide selected from the group consisting of: HIV RRE; HIV Tar; and BCR-AB1 fusion DNA sequences.
- A macromolecule of non-natural handedness of Claim 25 29. which binds to a target macromolecule and the target macromolecule is a phospholipid selected from the group consisting of: phosphoinositide; phosphoinositidase C; phosphoinositide 3-kinase; 10 phosphatidylinositol; phosphatidylinositol 3phosphate; phosphatidylinositol (4,5)bisphosphate; phosphatidylinositol (3,4,5)triphosphate; phosphatidylcholine; phosphatidylethanolamine; phosphatidic acid; inositol (1,4) bisphosphate; 15 inositol (1,4,5) triphosphate; diacylglycerol; sphingosine; sphingosine phosphate; sphigosine phosphocholine and ceramide.
- 30. A method of Claim 9 wherein the target macromolecule is vasopressin
 - 31. A method of Claim 30 where the L oligonucleotide that binds the target macromolecule is selected from the group consisting of SEQ ID NO.: 23, SEQ ID NO.: 234 and SEQ ID NO.: 25.
- 25 32. A process for producing a derivative of a macromolecule of non-natural handedness that binds to a target macromolecule of natural handedness, comprising the steps of:
- a) producing the macromolecule of non-natural

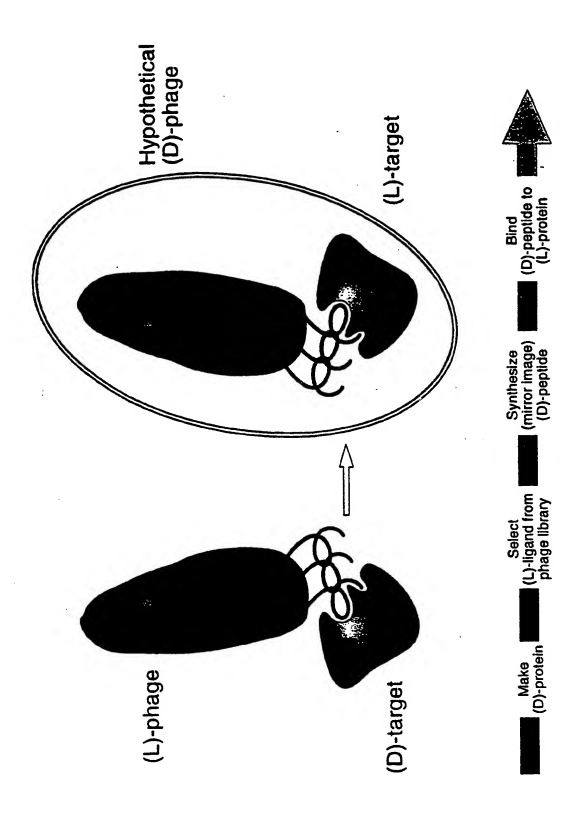
 handedness that binds to a target macromolecule
 of natural handedness;

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- b) providing an enantiomer of the target macromolecule or of a domain characteristic thereof;
- c) providing a library of macromolecule of natural handedness;
- d) contacting the library of c) with the enantiomer of b), under conditions appropriate for binding of a macromolecule of natural handedness in the library with the enantiomer of b), whereby the enantiomer of b) binds a macromolecule of natural handedness present in the library;
- e) producing the enantiomer of the macromolecule of natural handedness which is bound to the enantiomer of b), wherein the enantiomer of e) is a macromolecule of non-natural handedness which binds to the target macromolecule of natural handedness; and
- f) producing a derivative of the macromolecule of non-natural handedness of e).
- 20 33. A derivative obtainable by the process of Claim 20.



THE FIGURE

INTERNATIONAL SEARCH REPORT

Inter vial Application No PC1/US 96/06155

A. CLASSIE IPC 6	CO7K1/04 CO7H21/00 GO1N33/68	C07K14/00	
According to	International Patent Classification (IPC) or to both national classifica	tion and IPC	
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Minimum do	cumentation searched (classification system followed by classification	symbols)	
	on searched other than minimum documentation to the extent that suc		rehed .
Electronic de	ata base consulted during the international search (name of data base a	und, where practical, search terms used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the rele	vant passages	Relevant to claim No.
X	NATURE, vol. 368, 14 April 1994, LONDON G pages 651-653, XP002011492 P J FISHER ET AL.: "Calmodulin ir with amphiphilic peptides composed D-amino acids"	iteracts	18,21, 22,25, 27,33
	cited in the application see the whole document		10 21
X	NATURE, vol. 368, 21 April 1994, LONDON (pages 744-746, XP002011493 B J JAMESON ET AL.: "A rationally designed CD4 analogue inhibits experimental allergic encephalomy cited in the application see the whole document	,	18,21, 22,25, 27,33
		/	
X Fw	ther documents are listed in the continuation of box C.	Patent family members are listed i	n annex.
'A' docu	nent defining the general state of the art which is not	T later document published after the inte or priority date and not in conflict wil cited to understand the principle or th	h the application but
E' earlie	date nent which may throw doubts on priority claim(s) or	invention X' document of particular relevance; the cannot be considered novel or cannot involve an inventive step when the do Y' document of particular relevance; the	cument is taken alone
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'P' docu	r means ment published prior to the international filing date but than the priority date claimed	in the art. A document member of the same patent	
	e actual completion of the international search	Date of mailing of the international se	arch report
	23 August 1996		
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INTERNATIONAL SEARCH REPORT

Inter mal Application No PCI/US 96/06155

(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT Lategory * Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No.		
egory *	Cirian of document with numerous where appropriate of an enterior party	
······	SCIENCE, vol. 266, 23 December 1994, LANCASTER, PA US, pages 2019-2021, XP002011494	18,21, 22,25, 27,33
	C T DOOLEY ET AL.: "An all D-amino acid opioid peptide with central analgesic activity from a combinatorial library" cited in the application see the whole document	
	GENE, vol. 137, no. 1, 1993, AMSTERDAM NL, pages 13-16, XP002011495 K S LAM ET AL.: "Discovery of D-amino acid-containing ligands with selectide technology" see the whole document	18,21, 22,25, 27,32
	PEPTIDES, CHEMISTRY, STRUCTURE AND BIOLOGY. PROCEEDINGS OF THE 13TH AMERICAN PEPTIDE SYMPOSIUM, JUNE 20-25, 1993, EDMONTON, CANADA, 1994, ESCOM, LEIDEN, pages 984-985, XP002011496 C T DOOLEY ET AL.: "New, potent, N-acetylated all D-amino acid opioi"	18,21, 22,25, 27,32
	PEPTIDES, CHEMISTRY, STRUCTURE AND BIOLOGY. PROCEEDINGS OF THE 13TH AMERICAN PEPTIDE SYMPOSIUM, JUNE 20-25, EDMONTON, CANADA, 1994. ESCOM, LEIDEN, pages 1005-1006, XP002011497 K S LAM ET AL.: "Streptavidin-peptide interaction as a model system for molecular recognition " see the whole document	18,21, 22,25, 27,32
(PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF USA, vol. 92, no. 8, 11 April 1995, WASHINGTON US, pages 3110-3114, XP002011498 K ALEXANDROPOULOS ET AL.: "Proline-rich sequences that bind to SRC homology 3 domains with individual specificity" see the whole document	14
x	CELL, vol. 75, no. 1, 8 October 1993, NA US, pages 25-36, XP002011499 I GOUT ET AL.: "The GTPase dynamin binds to and is activated by a subset of SH3 domains " see the whole document	14
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INTERNATIONAL SEARCH REPORT

Inter 'onal Application No PC:/US 96/06155

		PC:/US 96/06155	
Continu:	(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	
х	CELL, vol. 76, no. 5, 11 April 1994, NA US, pages 933-945, XP002011500 H YU ET AL.: "Structural basis for the binding of proline-rich peptides to SH3 domains " see the whole document	14	
P , X	SCIENCE, vol. 271, 29 March 1996, LANCASTER, PA US, pages 1854-1856, XP002011501 T N M SCHUMACHER ET AL.: "Identification of D-peptide ligands through mirror-image phage display" see the whole document	1-32	
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